GNE.3230R1C39

PATENT -

IN THE

STATES PATENT AND TRADEMARK OFFICE

Applicant

Appl. No.

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For

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MEMBRANE

PTIDES AND NUCLEIC ENCODING THE SAME

Examiner

Blanchard

Group Art Unit

DECLARATIC

HRISTOPHER GRIMALDI, UNDER 37 C.F.R. § 1,132

Commissioner for Pate P.O. Box 1450 Alexandria, VA 22311

Dear Sir:

I, J. Christophe

, declare and say as follows:

- 1. I am a Genentech, Inc., South
- esearch Associate in the Molecular Biology Department of isco, CA 94080.
- 2. I joined Laboratory in the Mo numerous molecular b. analyses. I am current membrane associated cancer. In connection. directed the semi-qui Differential Tissue les specification that were: their normal counterpa-
- in January of 1999. From 1999 to 2003, I directed the Cloning ology Department. During this time I directed or performed niques including qualitative Polymerase Chain Reaction (PCR) ed in, among other projects, the isolation of genes coding for hich can be used as targets for antibody therapeutics against above-identified patent application, I personally performed or CR analyses in the assay entitled "Tumor Versus Normal Distribution" which is described in EXAMPLE 18 in the entify differences in gene expression between tumor tissue and
- 3. My scien and forms part of this
- culum Vitae, including my list of publications, is attached to (Exhibit A).
- 4. In diff differ significantly un
- expression studies, one looks for genes whose expression levels t conditions, for example, in normal versus diseased tissue.

Chromosomal aberra important markers o genes and their ence example, gene ampli duplicated, thus creati Gene under-expression chromosomal translo. each other chromosor: pattern relative to the [Singleton et al., Pat] [Grimaldi et al., Bloo: cancer cells a growth mechanism of tumor c aberration results in . polypeptide), as it de cancer therapy, for ear

- Compar important implicatio .3 this field are well aw a evidenced by an inc. over-expressed. him increased protein and gene is under-express. way, two cell sam; expected to have conta used to detect mkN... quantitative PCR, ... change in mRNA win these techniques was quantitation technical differentially expre : correspondingly single: expression is expression. decreased mRNA detection of increase treatment.
- of. How with the mRNA exand treatment. For with over- or und a identification of a classification and has

h as gene amplification, and chromosomal translocations are types of cancer and lead to the aberrant expression of specific eptides, including over-expression and under-expression. For s a process in which specific regions of a chromosome are le copies of certain genes that normally exist as a single copy. our when a gene is not transcribed into mRNA. In addition. cur when two different chromosomes break and are rejoined to ; in a chimeric chromosome which displays a different expression romosomes. Amplification of certain genes such as Her2/Neu 27Pt1:165-190], or chromosomal translocations such as t(5;14), 31-2085(1989); Meeker et al., Blood, 76(2):285-289(1990)] give il advantage relative to normal cells, and might also provide a ince to chemotherapy or radiotherapy. When the chromosomal expression of a mRNA and the corresponding gene product (the forementioned cases, the gene product is a promising target for he therapeutic antibody approach.

ene expression levels in normal versus diseased tissue has nostically and therapeutically. For example, those who work in the vast majority of cases, when a gene is over-expressed, as ction of mRNA, the gene product or polypeptide will also be tione identifies increased mRNA expression without associated is same principle applies to gene under-expression. When a product is also likely to be under-expressed. Stated in another ave differing mRNA concentrations for a specific gene are v different concentration of protein for that gene. Techniques Northern Blotting, Differential Display, in situ hybridization. fore recently Microarray technology all rely on the dogma that a similar change in protein. If this dogma did not hold true then atle value and not be so widely used. The use of mRNA lentified a seemingly endless number of genes which are tissues and these genes have subsequently been shown to have their protein levels. Thus, the detection of increased mRNA in increased polypeptide expression, and the detection of impected to result in decreased polypeptide expression. The polypeptide expression can be used for cancer diagnosis and

ill provides significant information useful for cancer diagnosis rer- or under-expression of a gene product does not correlate ... mRNA in certain tumor types but does so in others, then asion and protein expression enables more accurate tumor remination of suitable therapy. In addition, absence of over- or

е.

under-expression of t mRNA is crucial in. but the correspondi: will decide not to tr

7. I here that all statements restatements were made punishable by fine of Code and that such patent issued there.

By: Christo₁

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that in the presence of a particular over- or under-expression of the practicing clinician. For example, if a gene is over-expressed that is not significantly over-expressed, the clinician accordingly that agents that target that gene product.

tall statements made herein of my own knowledge are true and nation or belief are believed to be true, and further that these owledge that willful false statements and the like so made are a, or both, under Section 1001 of Title 18 of the United States neats may jeopardize the validity of the application or any

Date: 8 10 2004

GNE.3230R1C39

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

ព្រុវ្ម អ្នក Eaton, et al.

Appl. 10/063,557

: May 2, 2002

For : SECRETED AND

TRANSMEMBRANE

POLYPEPTIDES AND NUCLEIC

ACIDS ENCODING THE SAME

Examiner : David J. Blanchard

Group Art Unit: 1642

DECLARATION OF J. CHRISTOPHER GRIMALDI, UNDER 37 CFR §1.132

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Dear Sir:

- I, J. Christopher Grimaldi, declare and state as follows:
- 1. I am a Senior Research Associate in the Molecular Biology Department of Genentech, Inc., South San Francisco, CA 94080.
- 2. My scientific Curriculum Vitae, including my list of publications, is attached to and forms part of this Declaration (Exhibit A).
- I joined Genentech in January of 1999. From 1999 to 2003, I directed the Cloning Laboratory in the Molecular Biology Department. During this time I directed or performed numerous molecular biology techniques including semi-quantitative Polymerase Chain Reaction (PCR) analyses. I am currently involved, among other projects, in the isolation of genes coding for membrane associated proteins which can be used as targets for antibody therapeutics against cancer. In connection with the above-identified patent application, I personally performed or directed the semi-quantitative PCR gene expression analyses in the assay entitled "Tumor Versus Normal Differential Tissue Expression Distribution," which is described in EXAMPLE 18 in the specification. These studies were used to identify differences in gene expression between tumor tissue and their normal counterparts.
- 4. EXAMPLE 18 reports the results of the PCR analyses conducted as part of the investigating of several newly discovered DNA sequences. This process included developing

Appl. No. Filed

10/063,557 May 2, 2002

primers and analyzing expression of the DNA sequences of interest in normal and tumor tissues. The analyses were designed to determine whether a difference exists between gene expression in normal tissues as compared to tumor in the same tissue type.

- 5. The DNA libraries used in the gene expression studies were made from pooled samples of normal and of tumor tissues. Data from pooled samples is more likely to be accurate than data obtained from a sample from a single individual. That is, the detection of variations in gene expression is likely to represent a more generally relevant condition when pooled samples from normal tissues are compared with pooled samples from tumors in the same tissue type.
- 6. In differential gene expression studies, one looks for genes whose expression levels differ significantly under different conditions, for example, in normal versus diseased tissue. Thus, I conducted a semi-quantitative analysis of the expression of the DNA sequences of interest in normal versus tumor tissues. Expression levels were graded according to a scale of +, -, and +/- to indicate the amount of the specific signal detected. Using the widely accepted technique of PCR, it was determined whether the polynucleotides tested were more highly expressed, less expressed, or whether expression remained the same in tumor tissue as compared to its normal counterpart. Because this technique relies on the visual detection of ethidium bromide staining of PCR products on agarose gels, it is reasonable to assume that any detectable differences seen between two samples will represent at least a two fold difference in cDNA.
- 7. The results of the gene expression studies indicate that the genes of interest can be used to differentiate tumor from normal. The precise levels of gene expression are irrelevant; what matters is that there is a relative difference in expression between normal tissue and tumor tissue. The precise type of tumor is also irrelevant; again, the assay was designed to indicate whether a difference exists between normal tissue and tumor tissue of the same type. If a difference is detected, this indicates that the gene and its corresponding polypeptide and antibodies against the polypeptide are useful for diagnostic purposes, to screen samples to differentiate between normal and tumor. Additional studies can then be conducted if further information is desired.
- 8. I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information or belief are believed to be true, and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful statements may jeopardize the validity of the application or any patent issued thereon.

By: Christopher Grimoldi

Date:

te: 8 10/2004

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J. Christopher Grimaldi

1434-36th Ave. San Francisco, CA 94122 (415) 681-1639 (Home)

EDUCATION

University of California, Berkeley Bachelor of Arts in Molecular Biology, 1984

EMPLOYMENT EXPERIENCE

SRA

Genentech Inc., South San Francisco; 1/99 to present

Previously, was responsible to direct and manage the Cloning Lab. Currently focused on isolating cancer specific genes for the Tumor Antigen (TAP), and Secreted Tumor Protein (STOP) projects for the Oncology Department as well as Immunologically relevant genes for the Immunology Department. Directed a lab of 6 scientists focused on a company-wide team effort to identify and isolate secreted proteins for potential therapeutic use (SPDI). For the SPDI project my duties were, among other things, the critically important coordination of the cloning of thousands of putative genes, by developing a smooth process of communication between the Bioinformatics, Cloning, Sequencing, and Legal teams. Collaborated with several groups to discover novel genes through the Curagen project, a unique differential display methodology. Interacted extensively with the Legal team providing essential data needed for filing patents on novel genes discovered through the SPDI, TAP and Curagen projects. My group has developed, implemented and patented high throughput cloning methodologies that have proven to be essential for the isolation of hundreds of novel genes for the SPDI, TAP and Curagen projects as well as dozens of other smaller projects.

Scientist

DNAX Research Institute, Palo Alto; 9/91 to 1/99

Involved in multiple projects aimed at understanding novel genes discovered through bioinformatics studies and functional assays. Developed and patented a method for the specific depletion of eosinophils in vivo using monoclonal antibodies. Developed and implemented essential technical methodologies and provided strategic direction in the areas of expression, cloning, protein purification, general molecular biology, and monoclonal antibody production. Trained and supervised numerous technical staff.

Facilities

Manager Corixa. Rec

Corixa, Redwood City; 5/89 - 7/91.

Directed plant-related activities, which included expansion planning, maintenance, safety, purchasing, inventory control, shipping and receiving, and laboratory management. Designed and implemented the safety program. Also served as liaison to regulatory agencies at the local, state and federal level. Was in charge of property leases, leasehold improvements, etc. Negotiated vendor contracts and directed the purchasing department. Trained and supervised personnel to carry out the above-mentioned duties.

SRA

University of California, San Francisco Cancer Research Institute; 2/87-4/89.

Was responsible for numerous cloning projects including: studies of somatic hypermutation, studies of AIDS-associated lymphomas, and cloning of t(5;14), t(11;14), and t(8;14) translocations. Focused on the activation of hemopoietic growth factors involved in the t(5;14) translocation in leukemia patients...

Research Technician

Berlex Biosciences, South San Francisco; 7/85-2/87.

Worked on a subunit porcine vaccine directed against Mycoplasma hyopneumoniae. Was responsible for generating genomic libraries, screening with degenerate oligonucleotides, and characterizing and expressing clones in E. coli. Also constructed a general purpose expression vector for use by other scientific teams.

PUBLICATIONS

- Hilary F. Clark, et al. "The Secreted Protein Discovery Initiative (SPDI), a Large-scale Effort to Identify Novel Human Secreted and Transmembrane Proteins: a bioinformatics assessment." Genome Res. Vol 13(10), 2265-2270, 2003
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- 4. Jeanne Kahn, Fuad Mehraban, Gladdys Ingle, Xiaohua Xin, Juliet E. Bryant, Gordon Vehar, Jill Schoenfeld, J. Christopher Grimaldi (incorrectly named as "Grimaldi, CJ"), Franklin Peale, Apama Draksharapu, David A. Lewin, and Mary E. Gerritsen. "Gene Expression Profiling in an in Vitro Model of Angiogenesis." American Journal of Pathology Vol 156(6), 1887-1900, 2000.
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- Frances B. Lund, Nanette W. Solvason, Michael P. Cooke, Andrew W. Heath, J. Christopher Grimaldi, Troy D. Randall, R. M. E. Parkhouse, Christopher C Goodnow and Maureen C. Howard. "Signaling through murine CD38 is impaired in antigen receptor unresponsive B cells." European Journal of Immunology, Vol. 25(5), 1338-1345, 1995
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MANUSCRIPTS IN PREPARATION

 Sriram Balasubramanian, J. Christopher Grimaldi, J. Fernando Bazan, Gerard Zurawski and Maureen Howard. "Structural and functional characterization of CD38: Identification of active sité residues"

PATENTS

- 1. "Methods for Eosinophil Depletion with Antibody to CCR3 Receptor" (US 6,207,155 B1).
- "Amplification Based Cloning Method." (US 6,607,899)
- 3. Ashkenazi et al., "Secreted and Transmembrane Polypeptides and Nucleic Acids Encoding the Same." (this patent covers several hundred genes)
- 4. "IL-17 Homologous Polypeptides and Therapeutic Uses Thereof"
- "Method of Diagnosing and Treating Cartilaginous Disorders."

MEMBERSHIPS AND ACTIVITIES

Editor

Frontiers in Bioscience

Member

DNAX Safety Committee 1991-1999

Biological Safety Affairs Forum (BSAF) 1990-1991 Environmental Law Foundation (ELF) 1990-1991

The t(5;14) Chromosomal Translocation in a Case of Acute Lymphocytic Leukemia Joins the Interleukin-3 Gene to the Immunoglobulin Heavy Chain Gene

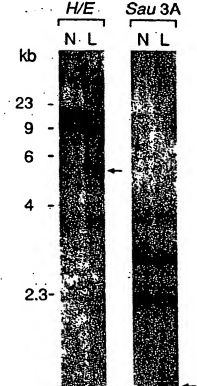
By J. Christopher Grimaldi and Timothy C. Meeker

Chromosomal translocations have proven to be important markers of the genetic abnormalities central to the pathogenesis of cancer. By cloning chromosomal breakpoints orne can identify activated proto-oncogenes. We have studled a case of B-lineage acute lymphocytic leukemia (ALL) that was essociated with peripheral blood sosinophilia. The canomosomal translocation t(6;14) (q31;q32) from this sample was cloned and studied at the molecular level. This

ing (Jh) region to the promotor region of the interleukin-3 (IL-3) gene in opposite transcriptional orientations. The data suggest that activation of the IL-3 gene by the enhancer of the immunoglobulin heavy chain gene may play a central role in the pathogenesis of this leukemia and the associated eosinophilla. • 1989 by Grune & Stratton, Inc.

translocation joined the immunoglobulin heavy chain join-

ARYOTYPIC STUDIES of leukemia and lymphoma have identified frequent nonrandom chromosomal translocations. Some of these translocations juxtapose the irmanunoglobulin heavy chain (IgH) gene with important protooncogenes, such as c-myc and bcl-2.13 In this way, the IgH gene can activate proto-oncogenes, resulting in disordered gene expression and a step in the development of cancer. The investigation of additional nonrandom translocations into the IgH locus allows us to identify new genes promoting the generation of leukemia and lymphoma. A distinct subtype of acute lymphocytic leukemia (ALL)



has been characterized by B-lineage phenotype, associated eosinophilia in the peripheral blood, and a t(5;14)(q31;q32) chromosomal translocation.14 This syndrome probably occurs in <1% of all patients with ALL. We hypothesized that the cloning of the translocation characteristic of this leukemia might allow the identification of an important gene on chromosome 5 that plays a role in the evolution of this disease. In this report we demonstrate that the interleukin-3 gene (IL-3) and the IgH gene are joined by this transloca-

MATERIALS AND METHODS Sample and DNA blots. A bone marrow aspirate from a representative patient with ALL (L1 morphology by French-American-British [FAB] criteria), peripheral eosinophilia (up to 20,000 per microliter with a normal value of <350 per microliter) and 2 t(5;14)(q31;q32) translocation was studied. Using published methods, genomic DNA was isolated and DNA blots were made. Briefly, 10 µg of high molecular weight (mol wt) DNA were digested using an appropriate restriction enzyme and electrophoresed on a 0.8% agarosé gel. The gel was stained with ethidium bromide, photo-

graphed, denatured, neutralized, and transferred to Hybond (Amorsham, Arlington Heights, IL). After treatment of the filter with ultraviolet light, hybridization was performed. The filter was washed to a final stringency of 0.2% saturated sodium citrate (SSC) and 0.1% sodium lauryly sulfate (SDS) and exposed to film. The human

From the Division of Hematology/Oncology, Department of

Jh probe has been previously reported.

Medicine, University of California, San Francisco. Submitted February 22, 1989; accepted March 8, 1989.

Supported by NIH Grant No. CA01102.

Address reprint requests to Timothy C. Meeker, MD, UCSF/ VAMC 111H, 4150 Clement St, San Francisco, CA 94121. Dr Grimaldi's current address is Biostan Inc. 440 Chesapeake

Genomic library. The genomic library was made using pub-

Dr, Seaport Centre, Redwood City, CA 94063.

The publication costs of this article were defrayed in part by page charge payment. This article must therefore be hereby marked "advertisement" In accordance with 18 U.S.C. section 1734 solely to indicate this fact.

o 1989 by Grune & Stratton, Inc. 0006-4971/89/7308-0031\$3.00/0

Fig 1. DNA blots of the leukemia sample. The restriction fregment pattern of normal human DNA (N) and the leukemis sample (L) were compared using a human Jh probe. Rearranged bands are indicated by arrows. Sample L exhibits a single rearranged band with both Hind III/EcoRI and SauSA restriction digests. The rearranged bends are less intense than the other bands because the majority of cells in the sample represent normal bone marrow elements.

1.3-

lished methods.³ Approximately 100 µg of high mol wt genomic DNA were partially digested with the Sau3A restriction enzyme. Fragments from 9 to 23 kilobases (kb) in size were isolated on a sucrose gradient and ligated into phage EMBL3A (Strategene, San Diego). Recombinant phage were packaged, plated, and screened as previously reported.⁵

DNA sequencing. Fragments for sequencing were cloned into M13 vectors and sequenced by the chain termination method using Sequenase (United States Blochemical, Cleveland). All sequence data were derived from both strands.

RESULTS

We studied a bone marrow sample from a patient with ALL and associated peripheral eosinophilia. Karyotypic analysis showed the characteristic t(5;14)(q31;q32) translocation. These features define a distinctive subtype of ALL.³⁴ The leukemic cells were analyzed for cell surface phenotype by immunofluorecence. They were positive for B1 (CD20), B4 (CD19), cALLA (CD10), HLA-DR, and terminal deoxynucleotidyl transferase (Tdt), but negative for surface immunoglobulin. This phenotypic profile describes an immature cell from the B-lymphocytic lineage.³

The leukemia DNA was analyzed by Southern blotting for rearrangements of the IgH gene. Using a human immuno-globulin Jh probe, a single rearranged band was detected by *EcoRI*, *HindIII*, *SstI*, *Sau3A*, and *EcoRI* plus *HindIII* restriction digests, suggesting rearrangement of one allele (Fig 1). The immunoglobulin Jh region from the other allele was presumably either deleted or in the germline configuration.

We hypothesized that the t(5;14)(q31;q32) juxtaposed a

growth-promoting gene on chromosome 5 with the immunoglobulin Jh region on chromosome 14. Therefore, a genomic
library was made from the leukemic sample and screened
with a Jh probe. Fifteen distinct positive clones were isolated
and screened for the presence of the rearranged Sau3A
fragment that was detected by DNA blotting. By this
analysis, five clones appeared to represent the rearranged
allele identified by DNA blots. One of these clones (clone no.
4) was chosen for further study and a detailed restriction
map was generated. The EcoRI, HindIII/EcoRI, and SstI
fragments from clone no. 4 that hybridized to the human Jh
probe were also identical in size to the rearranged fragments
from the leukemia sample, confirming that clone no. 4
represented the rearranged leukemic allele.

Phage clone no. 4 contained 3.7 kb of unknown origin joined to the IgH gene in the region of Jh4 (Fig 2). The IgH gene from Jh4 to the Cmu region appeared to be in germline configuration. Previously, the gene encoding hematopoietic growth factor IL-3 had been mapped to chromosome 5q31 so it was suspected that clone no. 4 might contain part of this gene. ⁵⁻¹² When the restriction map of human IL-3 and clone no. 4 were compared, they were identical for more than 3 kb (Fig 2).

We confirmed the juxtaposition of the IL-3 gene and the IgH gene by nucleic acid sequencing of the subcloned BstEII/HpaI fragment (Fig 2). The sequence of this fragment showed no disruption of the protein coding region or the messenger RNA of the IL-3 gene. The break in the IL-3 gene occurred in the promotor region, 452 base pairs (bp) upstream of the transcriptional start site (position 64, Fig

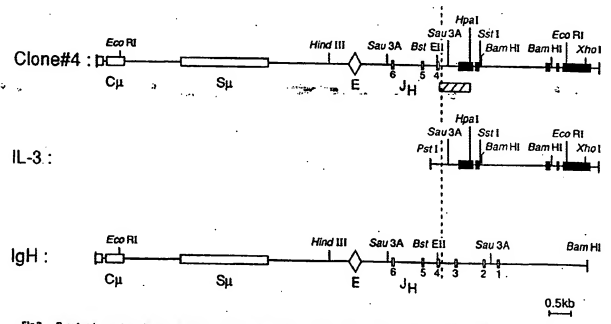


Fig 2. Brenkpoint region: t(5;14)(q31;q32). Comparative mapping of phage clone no. 4, the germline igJh region, and the germline iL-3 gene. The map of clone no. 4 is identical to that of igH until it diverges in the region of Jh4 (at the dashed line), after which it is identical to the map of iL-3. The two genes are positioned in a head-to-head orientation. The ig μ chain constant region (Cμ), switch region (Sμ), enhancer (E), and Jh segments are indicated (opan symbols). The five exons (dark boxes) and four introns of the IL-3 gene are shown. The

3A). The break in the IgH gene occurred 2 bp upstream of the Jh4 region. Between the two breaks, 25 bp of uncertain origin (putative N sequence) were inserted. Il. No sequences homologous to the immunoglobulin heptamer and nonamer could be identified in the IL-3 sequence (Fig 3B). Therefore, nucleic acid sequencing confirmed the juxtaposition of the IL-3 gene and the IgH gene. The sequence data clearly showed that the genes were positioned in opposite transcriptional orientations (head-to-head).

Available data also allowed us to determine the normal positions of the IL-3 gene and the GM-CSF gene in relation to the centromere of chromosome 5 (Fig 4). The IgH gene is kn own to be positioned with the variable regions toward the telemere on chromosome 14q. 215 It has also been shown that

GM-CSF maps within 9 kb of IL-3 in the same transcriptional orientation. 16 Using this information and assuming a simple translocation event in our sample, we can conclude that the IL-3 gene is normally more centromeric, and the GM-CSF gene more telomeric on chromosome 5q (Fig 4). Furthermore, both are transcribed with their 5' ends toward the centromere.

DISCUSSION

In this report we have cloned a unique chromosomal translocation that appears to be a consistent feature of a rare, yet distinct, clinical form of acute leukemia. This translocation joined the promotor of the IL-3 gene to the IgH gene. Except for the altered promotor, the IL-3 gene appeared

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Δ	5 GGTGACC	AGGGTTCCCTGGCCCAGTAGTCAAAGTAGTAGAGGTAATTCATCATAGCTGCGGATTAGCAGCGTGACCGGC TCCCAAGGGACCGGGGTCATCAGTTCATCATCTCCATTAAGTAGTATCGACGCCTAATCGTCGCACTGGCCG
H	3 CCACTGG	TCCCAGGGACCACAGATCAC
	5 TACCAGA 3 ATGGTCT	CAAACTCTCATCTGTTCGAGTGGCCTCCTGGCCACCCACC
	5 GTAGTCC	AGGTGATGGCAGATGAGATCCCACTGGGCAGGAGGCCTCAGTGAGCTGAGTCAGGCTTCCCCTTCCTGCCACA 248 TCCACTACCGTCTACTCTAGGGTGACCCGTCCTCCGGAGTCACTCGACTCAGTCCGAAGGGGGAAGGACGGTGT
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		•
	5 'CACA <u>TA'</u> 3 'GTGTAT	TAAGGCGGGAGGTTGTTGCCAACTCTTCAGAGCCCCACGAAGGACCAGAACAAGACAGAGTGCCTCCTGCCGAT 56LATTCCGCCCTCCAACAACGGTTGAGAAGTCTCGGGGTGCTTCCTGGTCTTGTTCTCTCACGAGGACGGCTA
	5 CCAAAC 3 GGTTTG	ATGAGCCGCCTGCCGTCCTGCTCCTGCTCCAACTCCTGGTCCGCCCCGGACTCCAAGCTCCCATGACCCAGAC TACTCGGCGGACGGCAGGACGACGACGAGGTTGAGGACCAGGCGGGGCCCTGAGGTTCGAGGGTACTGGGTTCTG
	5 AACGTC 3 TTGCAG	CTTGAGACAAGCTGGGTTAAC 3' 668 GGAACTTCTGTTCGACCCAATTG 5'
. 8	IgJh4	5 'TGGCCCCAGTAGTCAAAGTAGTCACATTGTGGGAGGCCCCATTAAGGGGTGCACAAAAAACCTGACTCTC 3 ' <u>ACCGGGGTCATCAGTTTCAT</u> CA <u>GTGTAAC</u> ACCCTCCGGGGTAATTCCCCACG <u>TGTTTTTGG</u> ACTGAGAG
		+++++++++++++++++++++++++
•	C1.#4	5 TGGCCCCAGTAGTCAAAGTAGTAGAGGTAATTCATCATAGCTGCGGATTAGCAGCGTGACCGGCTACCA 3 ACCGGGGTCATCAGTTTCATCA <u>TCTCCATTAAGTAGTATCGACGCCT</u> AATCGTCGCACTGGCCGATGGT

		•
	11-3	5 GCCACCAAGAGATGTGCTTCTCAGAGCCTGAGGCTGAACGTGGATGTTTAGCAGCGTGACCGGCTACCA 3 CCGTGGTTCTCTACACGAAGAGTCTCGGACTCCGACTTGCACCTACAAATCGTCGCACTGGCCGATGGT

Fig 3. Sequence of t(5;14)(q31;q32) breakpoint region. (A) Nucleotide sequence of the *BetEll/Hpal* tragment indicated on Fig 2. Nucleotides 1 to 36 represent the Jh4 coding region underlined on the coding strand. Nucleotides 39 to 63 are a putative N region. The sequence from position 64 to 668 is that of the germline R-3 gene. The R-3 TATA box (485), transcription start (516), and initiation metholine (567) are underlined. Two proposed regulatory sequences in the promotor are marked by asteriska (positions 182 and 389). (B) Comparative sequence of the t(6;14)(q31;q32) breakpoint region. The igJh4 region is shown with its coding region, haptamer, and transmer underlined. Clone no. 4 is shown with putative N region sequences underlined. The iL-3 sequence is also shown. A plus sign (+) donôtes the identified nucleotide between sequences. No haptamer or nonamer is identified in the tL-3 sequence.

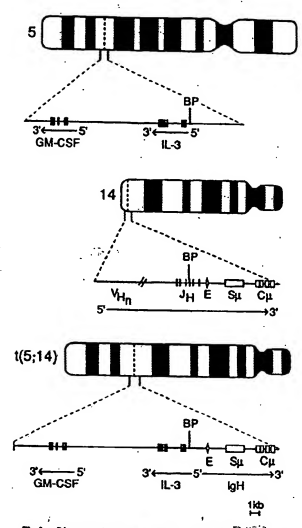


Fig 4. Diagram of the translocation. The normal chromosome 5q31 is shown with the GM-CSF gene telement to the fL-3 gene in the transcriptional orientation shown. On normal chromosome 14q32 the Vh regions are telement. The t[5;14][q31;q32] translocation results in the head-to-head orientation of these genes. Symbols are defined in Fig 2. BP, breakpoint position.

intact as no deletions, insertions, or point mutations were detected by restriction mapping of the entire gene and sequencing of part of the gene. The IgH gene has been truncated at the Jh4 region, which places the immunoglobulin enhancer within 2.5 kb of the IL-3 gene. This leads to the hypothesis that the enhancer is increasing transcription of a structurally normal IL-3 gene. The same mechanism is important for activation of the c-myc gene in some cases of Burkit's lymphoma. An alternate hypothesis is that the elimination of an upstream IL-3 promotor element is crucial to the activation of the IL-3 gene.

The proposed activation of the IL-3 gene suggests that an autocrine loop is important for the pathogenesis of this leukemia. Over-expression of the IL-3 gene coupled with

the presence of the IL-3 receptor in these cells could account for a strong stimulus for proliferation. In this regard, there are data indicating that immature B-lineage lymphocytes and B-lineage leukemias may express the IL-3 receptor. 21.22

An additional feature of this type of leukemia is the dramatic eosinophilia, consisting of mature forms. It has been hypothesized that the eosinophils do not arise from the malignant clone, but are stimulated by the tumor. 32,28 Because of the known effect of IL-3 on eosinophil differentiation, secretion of high levels of IL-3 by leukemic cells might have a role in the eosinophilia in this type of leukemia. 12

The data suggest that the recombination mechanism that is active in the IgH gene during normal differentiation has a role in this translocation. 13,14 This is supported by the breakpoint location at the 5' end of Jh4 and the presence of putative N-region sequences. On the other hand, no recombination signal sequence (heptamer and nonamer) was found in this region on chromosome 5, suggesting that additional factors also played a role. Further studies will elucidate the mechanism of this and other translocations.

In the leukemia we studied, it is possible that the immunoglobulin enhancer also activates the GM-CSF gene, since this gene is probably positioned only 14 kb away (Fig 4). This is known to be within the range of enhancer activation. The interleukin-5 (IL-5) gene maps to chromosome 5q31. Deregulation of the IL-5 gene by this translocation would act synergistically with IL-3 in the stimulation of eosinophil proliferation and differentiation. These and other questions will be answered by the study of more patient samples. We plan to determine whether the t(5;14)(q31;q32) translocation is capable of activating multiple lymphokines simultaneously and whether they cooperate in the generation of this leukemia.

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RAPID COMMUNICATION

Activation of the Interleukin-3 Gene by Chromosome Translocation in Acute Lymphocytic Leukemia With Eosinophilia

By Timothy C. Meeker, Dan Hardy, Cheryl Willman; Thomas Hogan, and John Abrams

The t(6;14)(q31;q32) translocation from B-lineage scute lymphocytic leukemia with easinophilis has been cloned from two leukemia samples. In both cases, this translocation joined the igH gene and the interleukin-3 (iL-3) gene. In one patient, excess IL-3 mRNA was produced by the leukemia cells. In the second patient, serum IL-3 levels were measured and shown to correlate with disease

A NUMBER OF chromosome translocations have been associated with human leukemia and lymphoma. In many cases the study of these translocations has led to the discovery or characterization of proto-oncogenes, such as bcl-2, c-abl, and c-myc, that are docated adjacent to the translocation.¹² It is now widely understood that cancer-associated translocations disrupt nearby proto-oncogenes.

A distinct subtype of acute leukemia is characterized by the triad of B-lineage immunophenotype, cosinophilia, and the t(5;14)(q31;q32) translocation. Leukemic cells from such patients have been positive for terminal deoxynucleotidyl transferase (Tdt), common acute lymphoblastic leukemia antigen (CALLA), and CD19, but negative for surface or cytoplasmic immunoglobulin. In previous work, we cloned the t(5;14) breakpoint from one leukemic sample (Case 1) and determined that the IgH and interleukin-3 (IL-3) genes were joined by this abnormality. In this report, we extend those findings by showing that the t(5;14)(q31;q32) translocation from a second leukemia sample (Case 2) has a similar structure, and we report our study of growth factor expression in these patients.

MATERIALS AND METHODS

Samples and Southern blots. Case 1 has been described. A Clinical features of Case 2 have been described in detail. DNA isolation and Southern blotting was done using previously described methods. Filters were hybridized with an immunoglobulin Jh probe, a 280 bp BamHI/EcoRI genomic IL-3 fragment, and an IL-3 cDNA probe. 13

Northern blots. RNA isolation and Northern blotting have been described. Briefly, Northern blots were done by separating 9µg total RNA on 1% agarose-formaldehyde gels. Equal RNA loading in each lane was confirmed by ethidium bromide staining. Blots were hybridized with an IL-3 cDNA probe extending to the Xho 1 site in exon 5, a 720 bp Sst I/Kpn I probe derived from intron 2 of the 1L-3 gene, a 600 bp Nhe 1/Hpa I IL-5 cDNA probe, and a 500 bp Pst I/Nco I granulocyte-macrophage colony stimulating factor (GM-CSF) cDNA probe.

Polymerase chain reaction. Primers were designed with BamHI sites for cloning. One primer hybridized to the Jh sequences from the IgH gene (Primer 144:5'-TAGGATCCGACGGTGACCAGGGT), and the other hybridized to the region of the TATA box in the IL-3 gene (Primer 161: 5'-AACAGGATCCCGCCTTATATGTGCAG). Polymerase chain reaction (PCR) (95°C for 1 minute, 61°C for 30 seconds, and 72°C for 3 minutes) was done using 500 ng genomic DNA and 50 pmol of each primer in 100 μL containing 67 mmol/L Tris-HCl pH 8.8, 6.7 mmol/L MgCl₂, 10% dimethyl sulfoxide (DMSO), 170 μg/mL bovine serum albumin (BSA) (fraction V),

activity. There was no evidence of excess granulocyte/macrophage colony stimulating factor (GM-CSF) or IL-5 expression. Our data support the formulation that this subtype of leukemia may arise in part because of a chromosome translocation that activates the IL-3 gene, resulting in autocrine and paracrine growth effects.

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16.6 mmol/L ammonium sulfate, 1.5 mmol/L each dNTP and Taq polymerase (Perkin-Elmer, Norwalk, CT). 13

Sequencing. Sequencing was done by chain termination in M13 vectors. As part of this study, we sequenced a subclone of a normal IL-3 promotor, covering 598 base pairs from a Sma I site at position—1240 (with respect to the proposed site of transcription initiation) to an Nhe I site at position—642. The plasmid containing this region was a gift from Naoko Arai of the DNAX Research Institute.

Expression in Cos7 cells. A genomic IL-3 fragment from Case 1 was cloned into the pXM expression vector. Briefly, the HindIII/Sal I fragment containing the IL-3 gene was subcloned from the previously described phage clone 4 into pUC18. The 2.6 kb fragment extending from the Sma I site 61 bp upstream of the IL-3 transcription start to the Sma I site in the polylinker was cloned into the blunted Xho I site of pXM. The negative control construct was the pXM vector without insert. Plasmids were introduced into Cos7 cells by electroporation, and supernatant was collected after 48 hours in culture.

TF1 bloassay. TF-1 cells were passaged in RPMI 1640 supplemented with 10% heat-inactivated fetal bovine serum, 2 mmol L-glutamine, and 1 ng/mL human GM-CSF. Il Samples and antibodies were diluted in this same medium lacking GM-CSF but containing penicillin and streptomycin. A 25 μ L volume of serial dilutions of patient serum was added to wells in a flat bottom 96-well microtiter plate. Rat anti-cytokine monocional antibody in a volume of 25 μ L was added to appropriate wells and preincubated for 1 hour at 37°C. Fifty microliters of twice washed TF-1 cells were added to each well, giving a final cell concentration of 1 \times 10° cells per well, (final volume, 100 μ L). The plate was incubated for 48 hours. The remaining cell viability was determined metabolically by the colori-

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© 1990 by The American Society of Hematology. 0006-4971/90/7602-0022\$3,00/0 Fig 1. Breakpoint sequences for Case 2. The germline IgJh5 region sequence (protein coding region and recombinations signal sequences are underlined) is on top, the translocation sequence from Case 2 (PCR primer sequences and putative N region are underlined) is in the middle, and the germline it-3 sequence, which we derived from a normal it-3 clone, is on the bottom.⁷ + indicates that each sequence has the same nucleotide. The sequence documents the head-to-head joining of the IL-3 and igH genes. The breakpoint in the it-3 gene occurred at position —834 (*).

metric method of Mosmann using a VMax microtiter plate reader (Molecular Devices, Menlo Park, CA) set at 570 and 650 nm. 16

Cytokine immunoassays. These assays used rat monoclonal anti-cytokine antibodies (10 µg/mL) to coat the wells of a PVC microtiter plate. The capture, antibodies used were BVD3-6G8, JES1-39D10, and BVD2-23B6, for the IL-3, IL-5, and GM-CSF assays, respectively. Patient sera were then added (undiluted and diluted 1:2 for IL-3, undiluted for IL-5, and undiluted and diluted 1:5 for GM-CSF). The detecting immunoreagents used were either mouse antiserum to IL-3 or nitroiodophenyl (NIP)-derivatized rat monoclonal antibodies JES1-5A2 and BVD2-21C11, specific for IL-5 and GM-CSF, respectively. Bound antibody was subsequently detected with immunoperoxidase conjugates: horseradish peroxidase (HRP)-labeled goat anti-mouse Ig for IL-3, or HRP-labeled rat (J4 MoAb) anti-NIP for IL-5 and GM-CSF. The chromogenic substrate was 3-3'azino-biz-benzthiazoline sulfonate (ABTS; Sigma, St Louis, MO). Unknown values were interpolated from standard curves prepared from dilutions of the recombinant factors using Softmax software available with the VMAX microplate reader (Molecular Devices).

RESULTS

Leukemic DNA from Case 2 was studied by Southern blotting. When digested with the HindIII restriction enzyme and hybridized with a human immunoglobulin heavy chain joining region (Jh) probe, a rearranged fragment at approximately 14 kb was detected (data not shown). When reprobed with either of two different IL-3 probes, a rearranged 14 kb

fragment, comigrating with the rearranged Jh fragment, was identified. When leukemic DNA was digested with HindIII plus BcoRI, a rearranged Jh fragment was detected at 6 kb. The IL-3 probes also identified a comigrating fragment of this size. These experiments indicated that the leukemic sample studied was clonal and that a single fragment contained both Jh and IL-3 sequences, suggesting a translocation had occurred.

To characterize better the joining of the IL-3 gene and the immunoglobulin heavy chain (IgH) gene, the polymerase chain reaction (PCR) was used to clone the translocation. A Jh primer and an IL-3 primer were designed to produce an amplified product in the event of a head-to-head translocation. While control DNA gave no PCR product, Case 2 DNA yielded a PCR-derived fragment of approximately 980 bp, which was cloned and sequenced.

The DNA sequence of the translocation clone from Case 2 confirmed the joining of the Jh region with the promotor of the IL-3 gene in a head-to-head configuration (Fig 1). Sequence analysis indicated that the breakpoint on chromosome 14 was just upstream of the Jh5 coding region. The breakpoint on chromosome 5 occurred 934 bp upstream of the putative site of transcription initiation of the IL-3 gene. We also determined that a putative N sequence of 17 bp was inserted between the chromosome 5 and chromosome 14 sequences during the translocation event. 17.11 Figure 2 shows

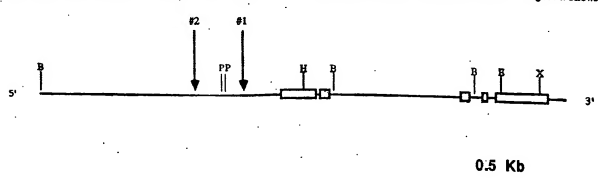


Fig 2. Relationship of chromosome 6 breakpoints to the IL-3 gane. This figure shows the two cloned breakpoints (arrows) in relation to the normal IL-3 gene. The breakpoint occurred at position —462 and the other at —834 (arrows). In both circumstances, the translocations resulted in a head-to-head joining of the IgH gene and the IL-3 gene, leaving the mRNA and protein coding regions of the IL-3 gene intact. Boxes denote the five IL-3 exons; restriction enzymes are [8] Bamill, (P) Pet I, (H) Host, (E) EcoRi, and (X) Xho I.

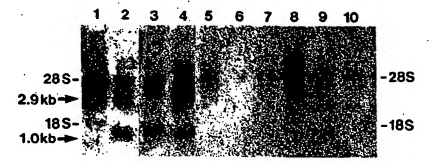


Fig. 3. Documentation of IL-3 mRNA over-expression. A Northern biot was prepared and hybridized with a probe for IL-3. Lane 1 contained RNA from unstimulated peripheral blood hymphocytes (PBL) as a negative control. Lane 2 contained RNA from PBL stimulated for 4hours with concenevalian A (ConA), and lane 3 contained RNA from PBL stimulated with ConA for 48 hours. As in the positive control lanes (2 and 3), a 1 to bend was identified in the leukemic sample from Case 1 (lane 4, lower arrow), suggesting aberrant expression of the IL-3 gene. In addition, the leukemic sample showed over-expression of an unspliced 2.9 kb IL-3 transcript (lane 4, upper arrow). We documented that this represented an unspliced precursor of the mature 1 kb transcript by showing that this band hybridized to a probe from intron 2 of the IL-3 gene. A similar 2.9 kb band was defected in nea 2, suggesting that an IL-3 mRNA of this size is cometimes districtable in normal mitogen-stimulated cells. Lane 5 through 10 represent RNA from six samples of B-lineage acute lymphocytic leukemia without the t(5):14) translocation, indicating that only the sample with the translocation exhibited IL-3 over-expression. Case 2 could not be analyzed by Northern blot because too few cells were available for study.

the locations of the two cloned breakpoints in relation to the JL-3 gene. The two chromosome 5 breakpoints were separated by less than 500 bp.

The genomic structure in Cases 1 and 2 suggested that a normal IL-3 gene product was over-expressed as a result of the altered promotor structure. This would predict that the IL-3 gene on the translocated chromosome was capable of making IL-3 protein. This prediction was tested by expressing a genomic fragment from the translocated allele of Case 1 containing all five IL-3 exons under the control of the SV40 promotor/enhancer in the Cos7 cell line. Cell supernatants were studied in a proliferation assay using the factor dependent crythroleukemic cell line, TF-1. The supernatants derived from transfections using the vector plus insert supported TF-1 proliferation, while supernatants from transfections using the vector alone were negative in this assay (data not shown). Furthermore, the biologic activity could be blocked by an antibody to human IL-3 (BVD3-6G8). This result showed that the translocated allele retained the ability to make IL-3 mRNA and protein.

The level of expression of IL-3 mRNA in leukemic cells from Case 1 was assessed. Northern blotting showed that the mature IL-3 mRNA (approximately 1 kb) and a 2.9 kb unspliced IL-3 mRNA were excessively produced by the leukemia (Fig 3). The 2.9 kb form of the mRNA is also present at low levels in normal peripheral blood T lymphocytes after mitogen activation (Fig 3). Several B-lineage acute leukemia samples without the t(5;14) translocation had undetectable levels of IL-3 mRNA in these experiments. In addition, although genes for GM-CSF and IL-5 map close to the IL-3 gene and might have been deregulated by the translocation, no IL-5 or GM-CSF mRNA could be detected in the leukemic sample (data not shown). 19,20

Three serum samples from Case 2 were assayed by immunoassay for levels of IL-3, GM-CSF, and IL-5 (Table 1). Serum IL-3 could be detected and correlated with the clinical course. When the patient's leukemic cell burden was

highest, the IL-3 level was highest. No serum GM-CSF or IL-5 could be detected.

Since the IL-3 immunoassay measured only immunoreactive factor, we confimed that biologically active IL-3 was present by using the TF-1 bioassay. This bioassay can be rendered monospecific using appropriate neutralizing monoclonal antibodies specific for IL-3, IL-5, or GM-CSF. We observed that sera from 1-16-84 and 3-14-84 contained TF-1 stimulating activity that could be blocked with anti-IL-3 MoAb (BVD3-6G8), but not with MoAbs to IL-5 (JES1-39D10) or GM-CSF (BVD2-23B6) (Fig 4; GM-CSF data not shown). The amount of neutralizable bioactivity in these two samples correlated very well with the difference in IL-3 levels obtained by immunoassay for these samples. Furthermore, the failure to block TF-1 proliferating activity with either anti-IL-5 or anti-GM-CSF was consistent with the inability to measure these factors by immunoassay and

Table 1. Peripheral Blood Counts and Growth Factor Levels at Different Times in Case 2

		Sample Date	_
	11/15/83	1/16/84	3/14/84
Peripheral blood counts (cells/µL)			
WBC	81,800	118,500	12,300
Lymphoblasts	0	33,785	. 0
Eosinophila	46,826	73,080	815
Serum growth factor levels (pg/mL)		•	
L-3	· <444	7,995	1,051
GM-CSF	<15	<15	<15
1L-6	<50	<60	<50

Peripheral blood counts from Case 2 at three different time points with the corresponding growth factor levels quantified by immunoessay. The patient received chemotherapy between 1/18/84 and 3/14/84 to lower his leukemic burden.³ No serum samples were evallable for a similar analysis of Case 1.

Abbreviation: WBC, white blood cells.

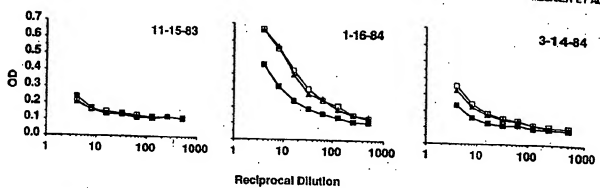


Fig. 4. Bloassay of serum R.-3. Leukemic patient sera were tested for bloactive it.-3 and it.-5 in the TF-1 proliferation assay. The reciprocal of the dilution is indicated on the horizontal axis and the optical density indicating the amount of proliferation is indicated on the vertical axis. Serum from all three time points was assayed eleutiteneously. The assay was rendered monospecific by using a 1 μg/ ml. final concentration of monoclonal rat anti-It.-3. BVD3-668 (EI), or anti-It.-5. JES1-39D10 (Δ): □ indicates no MoAb. On 1/16/64 and 3/14/84, inhibition of proliferation was evident in this presence of anti-It.-3 antibody, documenting serum levels of It.-3 on those days. Serum It.-6 was not detected in this assay, as anti-It.-6 did not after TF-1 proliferation.

indicated that these other myeloid growth factors were not detectably circulating in the serum of this patient.

DISCUSSION

In this report, we have extended our analysis of acute lymphocytic leukemia and eosinophilia associated with the t(5:14) translocation. In both cases we have studied, we have documented the joining of the IL-3 gene from chromosome 5 to the IgH gene from chromosome 14. The breakpoints on chromosome 5 are within 500 bp of each other, suggesting that additional breakpoints will be clustered in a small region of the IL-3 promotor. The PCR assay we have developed will be useful in the screening of additional clinical samples for this abnormality.

The finding of a disrupted IL-3 promotor associated with an otherwise normal IL-3 gene implied that this translocation might lead to the over-expression of a normal IL-3 gene product. In this work, we have documented that this is true. In addition, neither GM-CSF nor IL-5 are over-expressed by the leukemic cells. Furthermore, in one patient, serum IL-3 could be measured and correlated with disease activity. To our knowledge, this is the first measurement of human IL-3 in serum and its association with a disease process. The measurement of serum IL-3 in this and other clinical settings may now be indicated.

The finding of the IL-3 gene adjacent to a cancer-associated translocation breakpoint suggests that its activation is important for oncogenesis. It is our thesis that an autocrine loop for IL-3 is important for the evolution of this leukemia. The excessive IL-3 production that we have documented would be one feature of such an autocrine loop. The final proof of our thesis must await additional data. In particular, from the study of additional clinical samples, it will be necessary to document that the IL-3 receptor is present on the leukemic cells and that anti-IL-3 antibody decreases proliferation of the leukemia in vitro.

An important aspect of this work is the suggestion of a therapeutic approach for this disease. If an autocrine loop for IL-3 can be documented in this disease, attempts to lower circulating IL-3 levels or block the interaction of IL-3 with its receptor may prove useful. Because it is also possible that the cosinophilia in these patients is mediated by the paracrine effects of leukemia-derived IL-3, similar interventions may improve this aspect of the disease. Antibodies or engineered ligands to accomplish these goals may soon be available.

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Clinical and Pathologic Significance of the c-erbB-2 (HER-2/neu) Oncogene

Timothy P. Singleton and John G. Strickler

The c-srbB-2 oncogene was first shown to have clinical significance in 1987 by Slamon et al, who reported that c-srbB-2 DNA amplification in breast carcinomas correlated with decreased survival in patients with metastasis to axillary lymph nodes. Subsequent studies, however, of c-srbB-2 activation in breast carcinoma reached conflicting conclusions about its clinical significance. This oncogene also has been reported to have clinical and pathologic implications in other neoplasms. Our review summarizes these various studies and examines the clinical relevance of c-srbB-2 activation, which has not been emphasized in recent reviews. 57.34.55 The molecular biology of the c-srbB-2 oncogene has been extensively reviewed. 37.32.55 and will be discussed only briefly here.

BACKGROUND

The c-erbB-2 oncogene was discovered in the 1980s by three lines of investigation. The new oncogene was detected as a mutated transforming gene in neuroblastomas induced by ethylnitrosurea treatment of fetal rats. A.TA.TA.T. The c-erbB-2 was a human gene discovered by its homology to the retroviral gene v-erbB. A.S.TA.TA.T. HER-2 was isolated by screening a human genomic DNA library for homology with v-erbB. When the DNA sequences were determined subsequently, c-erbB-2, HER-2, and new were found to represent the same gene. Recently, the c-erbB-2 oncogene also has been referred to as NGL.

The c-erbB-2 DNA is located on human chromosome 17q2124.364 and codes for c-erbB-2 mRNA (4.6 kb), which translates c-erbB-2 protein (p185). This

166 T.P. SINGLETON AND J.G. STRICKLER

protein is a normal component of cytoplasmic membranes. The e-erbB-2 oncogene is homologous with, but not identical to, e-erbB-1, which is located on chromosome 7 and codes for the epidermal growth factor receptor. The e-erbB-2 protein is a receptor on cell membranes and has intracellular tyrosine kinase activity and an extracellular binding domain. Electron microscopy with a polyclonal antibody detects e-erbB-2 immunoreactivity on cytoplasmic membranes of neoplasms, especially on microvilli and the non-villous outer cell membrane. In normal cells, immunohistochemical reactivity for e-erbB-2 is frequently present at the basolateral membrane or the cytoplasmic membrane's brush border. ***Estate: ***Estate:

There is experimental evidence that o-erbB-2 protein may be involved in the pathogenesis of breast neoplasia. Overproduction of otherwise normal c-erbB-2 protein can transform a cell line into a malignant phenotype. Also, when the neu oncogene containing an activating point mutation is placed in transgenic mice with a strong promoter for increased expression, the mice develop multiple independent mammary adenocarcinomas. In other experiments, monoclonal antibodies against the neu protein inhibit the growth (in nude mice) of a neu-transformed cell line, the maximum and immunization of mice with neu protein protects them from subsequent tumor challenge with the neutransformed cell line. Some authors have speculated that the use of antagonists for the unknown ligand could be useful in future chemotherapy. Further review of this experimental evidence is beyond the scope of this article.

The c-erbB-2 activation most likely occurs at an early stage of neoplastic development. This hypothesis is supported by the presence of c-erbB-2 activation in both in situ and invasive breast carcinomas. In addition, studies of metastatic breast carcinomas usually demonstrate uniform c-erbB-2 activation at multiple sites in the same patient, 11.12.36.13.26 although c-erbB-2 activation has rarely been detected in metastatic lesions but not in the primary tumor. 57.68.107 Even more rarely, c-erbB-2 DNA amplification has been detected in a primary breast carcinoma but not in its lymph node metastasis. In patients who have bilateral breast neoplasms, both lesions have similar patterns of c-erbB-2 activation, but only a few such cases have been studied. 11

MECHANISMS OF c-erbB-2 ACTIVATION

The most common mechanism of c-erbB-2 activation is genomic DNA amplification, which almost always results in overproduction of c-erbB-2 mRNA and protein. 17,24,65,81 The c-erbB-2 amplification may stabilize the overproduction of mRNA or protein through unknown mechanisms. Human breast carcinomas with c-erbB-2 amplification contain 2 to 40 times more c-erbB-2 DNA45 and 4 to 128 times more c-erbB-2 mRNA5450 than found in normal tissuc. Most human breast carcinomas with c-erbB-2 amplification have 2 to 15 times more c-erbB-2 DNA. Tumors with greater amplification tend to have greater overproduction. 17,53,65 The non-mammary neoplasms that have been studied tend to have

167

similar levels of c-erbB-2 amplification or overproduction relative to the corresponding normal tissue.

The second most common mechanism of c-erbB-2 activation is overproduction of c-erbB-2 mRNA and protein without amplification of c-erbB-2 DNA.61 The quantities of mRNA and protein usually are less than those in amplified cases and may approach the small quantities present in normal breast or other tissues. 17.50.52 The c-erbB-2 protein overproduction without mRNA overproduction or DNA amplification has been described in a few human breast carcinoma cell lines.47

Other rare mechanisms of c-erbB-2 activation have been reported. Translocations involving the c-erbB-2 gene have been described in a few mammary and gastric carcinomas, although some reported cases may represent restriction fragment length polymorphisms or incomplete restriction enzyme digestions that mimic translocations. 31.65,74.450,108 A single point mutation in the transmembrane portion of neu has been described in rat neuroblastomas induced by ethylnitrosurea. 2.55 The mutated neu protein has increased tyrosine kinase activity and aggregates at the cell membrane. 10,20,20 Although there has been speculation that some of the amplified c-erbB-2 genes may contain point mutations,4 none has been detected in primary human neoplasms. 41,50,61

TECHNIQUES FOR DETECTING 6-orbB-2 ACTIVATION

Detection of c-erbB-2 DNA Amplification

Amplification of c-erbB-2 DNA is usually detected by DNA dot blot or Southern blot hybridization. In the dot blot method, the extracted DNA is placed directly on a nylon membrane and hybridized with a c-erbB-2 DNA probe. In the Southern blot method, the extracted DNA is treated with a restriction enzyme, and the fragments are separated by electrophoresis, transferred to a nylon membrane, and hybridized with a c-erbB-2 DNA probe. In both techniques, c-erbB-2 amplification is quantified by comparing the intensity (measured by densitometry) of the hybridization bands from the sample with those from control tissue.

Several technical problems may complicate the measurement of c-erbB-2 DNA amplification. First, the extracted tumor DNA may be excessively degraded or diluted by DNA from stromal cells. 53 Second, the c-erbB-2 DNA probe must be carefully chosen and labeled. For example, oligonucleotide cerbB-2 probes may not be sensitive enough for measuring a low level of c-erbB-2 amplification, because diploid copy numbers can be difficult to detect (unpublished data). Third, the total amounts of DNA in the sample and control tissue must be compensated for, often with a probe to an unamplified gene. Many studies have used control probes to genes on chromosome 17, the location of cerbB-2, to correct for possible alterations in chromosome number. Identical results, however, are obtained by using control prohes to genes on other chromosomes, s. ic. with rare exception. 17 Studies using control probes to the beta-

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globin gene must be interpreted with caution, because one allele of this gene is deleted occasionally in breast carcinomas.

Amplification of c-erbB-2 DNA was assessed by using the polymerase chain reaction (PCR) in one recent study. Oligoprimers for the c-erbB-2 gene and a control gene are added to the sample's DNA, and PCR is performed. If the sample contains more copies of c-erbB-2 DNA than of the control gene, the c-erbB-2 DNA is replicated preferentially.

Detection of c-erbB-2 mRNA Overproduction

Overproduction of c-erbB-2 mRNA usually is measured by RNA dot blot or Northern blot hybridization. Both techniques require extraction of RNA but otherwise are analogous to DNA dot blot and Southern blot hybridization. Use of PCR for detection of c-erbB-2 mRNA-har-been described in two recent abstracts. 4.100

Overproduction of c-erbB-2 mRNA can be measured by in situ hybridization. Sections are mounted on glass slides, treated with protease, hybridized with a radiolabeled probe, washed, treated with nuclease to remove unbound probe, and developed for autoradiography. Silver grains are seen only over tumor cells that overproduce c-erbB-2 mRNA. Negative control probes are used. ALEM OUT experience indicates that these techniques are relatively insensitive for detecting c-erbB-2 mRNA overproduction in routinely processed tissue. Although the sensitivity may be increased by modifications that allow simultaneous detection of c-erbB-2 DNA and mRNA, in situ hybridization still is cumbersome and expensive (unpublished data).

All of the above c-erbB-2 mRNA detection techniques have several problems that make them more difficult to perform than techniques for detecting DNA amplification. One major problem is the rapid degradation of RNA in tissue that is not immediately frozen or fixed. In addition, during the detection procedure, RNA can be degraded by RNase, a ubiquitous enzyme, which must be eliminated meticulously from laboratory solutions. Third, control probes to genes that are uniformly expressed in the tissue of interest need to be carefully selected.

Detection of o-erbB-2 Protein Overproduction

The most accurate method is for detecting c-erbB-2 protein overproduction are the Western blot method and immunoprecipitation. Both techniques can document the binding specificity of various antibodies against c-erbB-2 protein. In Western blot studies, protein is extracted from the tissue, separated by electrophoresis (according to size, transferred to a membrane, and detected by using antibodies to c-erbB-2. In immunoprecipitation studies, antibodies against c-erbB-2 are added to a tumor lysate, and the resulting protein-antibody precipitate is separated by gel electrophoresis and stained for protein. Both Western blot and immunoprecipitation are secful research tools but currently are not practical for diagnostic pathology. Two recent abstracts have described an enzyme-linked immunosorbent assay (E - A) for detection of c-erbB-2 protein. [8.5]

some antibodies detect cin fixed tissue. In gene diagnose c-erbB-2 protei. antibody to antibody. De it can be detected in in **without fixation or** freezi.

Overproduction of e-mini-2 protein is most commonly assessed by various immunohistochemical techniques. These procedures often generate conflicting results, which are explain an least partially by three factors. First, various studies have used differe it polyclonal and monoclonal antibodies. Because some polyclonal antibodies recognize weak bands in addition to the c-erbB-2 protein band on Western blot or immunoprecipitation, the results of these studies should be interpresed with caution. \$2,75,47,61 Even some monoclonal antibodies immunoprecipitate protein bands in addition to c-erbB-2 (p185). 20,50,000 Second, tissue fixation continues to variability between studies. For example, 1-2 protein only in frozen tissue and do not react brinalin fixation diminishes the sensitivity of immunohistochemical me e s and decreases the number of reactive cells.41,50 When Bouin's fixative is there may be a higher percentage of positive cases. 22 Third, minimal criteria for interpreting immunohistochemical staining are generally lacking. Alt note the there is general agreement that distinct crisp cytoplasmic membrane s. ...ing is diagnostic for c-erbB-2 activation in breast carcinoma, the number our vitive cells and the staining intensity required to . production varies from study to study and from on of c-erbB-2 protein is not a problem because arm more than 24 hours after tumor resection

ACTIVATION OF c-erb.

incidence of c-erbB-2 Most studies of c-erbBtypes of infiltrating breas in 19.1 percent (519 of 2 c-erbB-2 mRNA or prote 2714) of invasive carcino:

erbB-2 mRNA or protein t that lacked c-erbB-2 DN The incidence of c-c

with the histological sub ing ductal carcinomas data. Other variants of inflammatory carcinoma 11), and medullary care: tion is infrequent in it tubular carcinoma (7 pc

The c-erbB-2 prote. ductal carcinomas in sit 49 of 72). The micropar c-erbB-2 activation, 40.54.

REAST LESIONS

rene activation do not specify histological suboma, Amplification of c-erbB-2 DNA was foundvasive carcinomas in 25 studies (Table 1), and regoduction was detected in 20.9 percent (568 of in 20 studies. Twelve studies have documented creduction in 15 percent (88 of 604) of carcinomas ..ication.

ctivation in infiltrating breast carcinoma varies proximately 22 percent (142 of 650) of infiltrat-,53-2 activation, as expected from the above cinoma with frequent c-erbB-2 activation are . ent, 54 of 87), Paget's disease (82 percent, 9 of : percent, 5 of 23). In contrast, c-erbB-2 activan; lobular carcinoma (7 percent, 5 of 73) and

.oduction is present in 44 percent (44 of 100) of scially comedocarcinoma in situ (68 percent, · · · if ductal carcinoma in situ also tends to have ile if larger cells are present. The greater fre-

	G-erbB-2 DNA	o-erbB-2 mRNA	o-exten-2 Protein
Histological Diagnosis	Amptification*	Qwerproduction	Overproduction
		•	:
	19/103 78 15/95.20	,	24/53, sz +, 'D
	15,86 11 17,73,7	٠ - د	22/45, 11/36, H
•	16/56,4 6/61,8		7/24,*** 1/1081
	11/57,2 10/57,45	•	
	13/51,13 6/48,19		
5	10/38,12/38,1	•	,
	1/25,13 7/24,11		
- 0	7/15,31 7/10,1	•	
		::3	
Carchoma, type not specified but lacking c-erbB-2 DNA emplification	1	16/136,** 14/73,** BAS,** QB,** 1/4,** Q3**	16/231, 770 18/138, 77 13/35, 13/14/23, 23 1/28, # 3/24, 44 0/17**
Inditraling ductal carchoma	21/118, ⁴² 23/107, ³⁴ 17/50, ⁴⁴ 7/57 ⁴⁰	35/85*	22/137,40 14/83,40 8/34/40
1. 3	14/53 (comedo- carchroma) ¹⁴ 3/53 (ubuloduotal		

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Manustra carrinoma	33/80's 3/62	46775	200
Donale demons	•	ı	546. = 2/3. th 2/2 ^{cd}
Total day commonts	0/6.4 0/1 ²⁰		1/840
Medulary carcinoma	2/4,11 0/1*		1/12.to 1/3,to 1/2,to 0/13
Mucinous carchoma	071,* 071 ^{ss}	1 1 114	1/2**
Inflitrating tobular carchoma	1/15,10 0/674	- 1 79.	2/27,¤ 0/12,* 0/8,** 1/5**
Warren and Andrews	0,780		
		<u>:</u>	3
ייין ואיסויי	, i		2000 to 2000 t
Cuctal CIS, solid or comedo type	١.	!	10/105
	1	.:	10/100
Ourtai Cis, meropaphiany type	ı		1(local)/14**
Ductal CIS, micropapillary or crumorm type	- 1		0/16,1 1/9, 1 0/30
Ductal CIS, papillary or enforcem type	1		. 0/16~
- Chuler CIS	}		

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own as number of cases with activatorynumber of cases studied; informaties to govern as a superass protein studies used Western blots; the rest used immunohistochemical methods. quency of c-erbB-2 protein a pared with infiltrating ductal care many infiltrating ductal care noma, which show c-erbB-2 that careinoma in situ with srbB-2 activation during procomponents of ductal carein erbB-2 activation, 11,29 although the immunohistochemical s noma. Activation of celesions contain more than erbB-2 protein overproduct but may include other are erbB-2 protein in ductal careibB-2 protein in ductal car

Activation of c-erb 8-2 including fibrocystic diseas, membrane immunohistoche in atypical ductal hyperplas has been noted infrequer i diploid, and c-erb B-2 is ex;

These preliminary da for resolving many of the coexample, c-erbB-2 activat: In addition, because c-erb1 cribriform carcinoma in sit; 2 activation in these lesions histological features of coicc-erbB-2, are unlikely to

aduction in comedocarcinoma in situ, commona, could be explained by the fact that arise from other types of intraductal carcitation infrequently. Others have speculated 2-2 activation tends to regress or to lose conto invasion. **Makes** Infiltrating and in situ cover, usually are similar with respect to cauthors have noted more heterogeneity of mattern in invasive than in in situ carcitis infrequent in lobular carcinoma in situ. If the control of carcinoma in situ, the control of coccur in the comedocarcinoma in situ reinoma in situ. ***Comedocarcinoma in situ correlates with larger cell size and a

been identified in benign breast lesions, all momas, and radial scars (Table 2). Strong divity for e-erbB-2 has not been described over taccentuation of membrane staining normal breast tissue, e-erbB-2 DNA is wer levels than in activated tumors. M.S. S. Lace erbB-2 activation may not be useful thems in diagnostic surgical pathology. For ent in tubular careinoma and radial scars. It is unusual in atypical ductal hyperplasia, we reinoma in situ, detection of e-erbB-cally lid in their differential diagnosis. The

TABLE 2. o-orbB-2 ACTIVAT.

Histological Diagnosis
Fibrocyetic disease
Atypical ductal hyperplasia
Benign ductal hyperplasia
Scierosing adenosis
Fibroadenomas
Radial scare
Blunt duct adenosis
"Bresst mastosis"
Shown as number of cases with a

111 ... JHEAST LÉSIONS

_	DD-2 mRNA	o-erbB-2 Protein Overproduction
	-	0/32,30/9,40/850
		2(weak)/21, ⁵⁴ 1(cytoplasmio)/13 ³⁰
		0/1233
		0/439
	^{.1} . C\ 234	0/21, ³⁰ 0/10, ⁴⁰ 0/8, ⁴⁰ 0/3 ⁻²
	-	0/22**
	_	0/1439
5	a	_ `

I, reference la given as a superscript

tr.

c-erbB-2, however, doe lobular carcinoma. Furt!

Correlation of c-erbB-Multiple studies have at pathologic prognostic faction with lymph node metast of 17 series, and with his associated with e-crbB-2 content and low prolife prognostic factors and in

Correlation of c-erhib-Various studies have atte features that may pred correlated with absence sence of progesterone rdid not correlate with a erhB-2 activation was a

Correlation of c-orbid-Slamon et al^{70,01} first sin pendently predicts decorrelation of c-orbidthe correlation of mainiet al also reported the tor only in patients.

A large number. Description with province tween c-erbB-2 are a soft these series, the relation of the correlation of explanations for the series.

One problem mainly in patient: studies of patients with c-srbB-2 activatio without axillary coutcome. Table (cout axillary met studies with a himbetween c-srbB-evidence suggest with metastasis to

etal carcinoma over infiltrating tea would be useful.

thologic Prognostic Factors 2-crbB-2 activation with various don of c-erbB-2 was correlated th higher histological grade in 6 ries. Large tumor size was not is (11 of 14). Tetraploid DNA 33-67, have been suggested as 2 activation.6.7

col-Prognostic Factors
-erbB-2 activation with clinical
de 4). Activation of c-erbB-2
10 of 28 series and with abin most studies, patient age
in the rest of the reports, cger or older ages.

Int Outcome

the c-erbB-2 oncogene indeswith breast carcinoma. The decome was nearly as strong asis with poor outcome. Slamon important prognostic indica-

s, there was a correlation beer decreased survival. In five retivation was reported to be ast, 18 series did not confirm once or survival. Four possible low.

n correlates with prognosis summerized in Table 5, most istasis showed a correlation of last, most studies of patients of a correlation with patient in all attents (with and withgroup. There is a trend for lases co-show an association. Thus, most of the current gnostic value only in patients

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Prognostic Factor

30.0°

Overproduction

o-erbB-2 Protein Overproductions

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. (176)101 (168)11 (38)13 13501 ~ (186)~ (137) (86) (63)" (58)11 (50)4 (41) (47)13 (17, رد) (درا)ت 43 7 Higher

"A correlation to statis ically significant at <0.05, equivocal at best between 0.05 and 0.15, and not statistically elgosticant at >0.15.

Pklumbers hade parenthases are the number of patients in an individual study, asperangly is the reference. Some studies-graphized more than one group of patients.

Spy Western blot method, all other protein studies used immanoristanchemical methods.

TABLEA

Prognostic Factor P : A	cerbB-2 DNA	o-erbB-2 mRNA	cerb8-2 Protein
	Amplification	Overproduction	Overproduction

(62) אַיַּבְּטָרָ**) אַנְאַבְּאַ (**(123) פּייַּ(124) אַנְאָבָאַ (125) פּייַּ(124) אָנְאָבָאָן אָרִרּרָן) (55)2 (53)11 (45)12 (56)22 (53)11 (49)13 (41)22 (15)23 7.0.16

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*A correlation is statisficatly etynificant at <0.05, equivocal at best between 0.05 and 0.15, and not statisficably eignificant at >0.15

Phumbers trade parentheses are the number of patients in an individual study; superscript is the reference. Some studies analyzed more than one group of patients

The reference is the method; all other protein studies used immunicitisational methods.

176 T.P. SINGL

TABLE & CORREL WITH BREAST CAR

	Type c
	e-erb!
P1	Activ
< 0.05	DNA
<0.05	DNA
<0.05	DNA
< 0.05	DNA
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< 0.05	DNA
<0.05	DNA
< 0.05	DNA
< 0.05	DNA
< 0.05	Protei +
< 0.05	Protein
0.05-0.15	DNA
0.05-0.15	Protein
0.05~0.15	Proto 1
>0.15	DNA
>0.15	DNA
>0.15	DNA
>0.15	mRN.
>0.15	Prote .
>0.15	Pro tein
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>0.15	Proteh
>0.15	Prote
>0.15	Prote .
>0.15	Protein

The endpoints of the emb8-2 activation and at 0.05 to 0.15, and is Shown as variable in immunohistochemical at multivariate stati-

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HOUTCO HE IN PATIENTS

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	. м	81	•
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41 .	ر .	86	•
41	IJ	40	

d survival or both. Correlation between o-

em blot; the other protein studies used

177

TABLE 6. PERCE WITH PROGNOS

% of tumors with lymph node metastasis in each study

P for correlation of c-Each study's percent erbB-2 activation are carbor patients, who are the types of c-erc

A second; together in man; c-erbB-2 active infiltrating duc erbB-2 activate inflammatory berbB-2 activationoma, but it is a

A third potci-erbB-2 activa metastasis. Two metastasis who and absence of recurrence. 22,67 no association 1.

A fourth p correlates better overproduction

ATTACKE!	TASTA SIS	COMPARED
ATION		

64(DNA)***	64 (mRNA)∞ 61(DNA)⁴
, m	58 (DNA)** 57 (DNA)** 55 (Protein)**
	48(Protein) ¹¹ 48(Protein) ¹²
42(Protein)**	
	i
P<0.15	P>0.15

empared with the correlation between othat considered, as one group, all breast cripts are the references. In parentheses

reast carcinoma are grouped creent literature suggests that inoma, studies that combine ite the prognostic effect of emost st dies do not analyze andition frequently shows can the usual mammary carci-

dies that attempt to correlate s of bres st carcinoma without patients without lymph node nee (such as large tumor size rexpression predicted early situ, sas small study found oB-2 activation.

ing whether the prognosis of or with mRNA or protein a between c-erbB-2 activa-

and breast carcinoma pa de constant para de constant carcinoma pa de constant poor resurvival. The constant poor resurvival po

Comparison of c-erbB : / · · v Breast Carcinoma

Other oncogenes that me are reviewed elsewhere between the clinical rele

The c-myc gene is c tion generally has less pr One study found a corremyc, although other representations of however, could demonst a more prognostic importa

The gene c-erbB-1 homologous with c-erbB Overproduction of ECF and may correlate with a erbB-2 and ECFR in the poor prognostic factors. 30 amplification of c-erbB-2 although at the molecul protein. 51,82,61,84,100 Recei

The genes c-erbA at tor, and they are located are frequently coamplific-erbA expression in bre tant role for this gene in without ear-1 amplification similar to tumors with b c-erbB-2 amplification so or ear-1.

Other genes also h. carcinomas. One study & 2 mRNA and increased and Ki-ras. 105 Allelic de breast carcinoma. 11 but i studies have suggested . breast carcinoma and ac

A amplification (Table 5), ion of e-erbB-2 may have tation has more prognossignificance of e-erbB-2 arther esearch. 17.22 Few e with e-erbB-2 mRNA overproduction use relamical studies with poly-

acogenes in

in hus an breast cancertricted to a comparison ther or rogenes.

omas, out c-myc activaoB-2 activation. u.s.m.s. NAs o. c-erbB-2 and ciso Sub equent research, mas in which c-myc has

tor re .ptor (EGFR) is in b. st carcinomas. 79 ently an amplification have xamined both ctronge correlation with no co relation between 1 of c-c B-2 and EGFR. phory: tion of c-erbB-2 east e einomu. 4100 normone recepbyro osom .7.:These genes aom The absence of ience gainst an imporn of c :rbB-2 can occur cerea. I survival that is catio.. Consequently, cation of c-erbA : amp

3-2 a ration in breast ween preased c-erbBgrove factor chain A, er prognosis in rbBstag r recurrence of coger 12.13

ACTIVATION OF C .

Incidence of c-er.
Table 7 summarized detected, usually w.

TABLE 7. PRESENCE NORMAL HUMAN TIS

Thisues With o-erbB-2 mRNA	Tiss c-
Skina	Eplot
	Exter
	Eccrl
•	Fetal
	Foto
. Stomach ²⁴	Stor
	Fetal
Jejunum³4	Smal
Colones	Colu.
Kidney ²⁴	Feta
•	Fetal
	Dista
	Feli
	Feli
	Feli
Liver ²⁴	Hep.
	. Pari
	Pai
	End of
Lamott	
Lung** }	Fetu Fet
	Brc
Patal hardany	ac
Fetal brains	Feta
The control	reta
Thyrold ¹	
Uterus ^e	_
	Ov.
	Bloc

This protein study use.

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nmary 116 sues c-erbB-2 expression has been tethods using polyclonal anti-

A OR See aB-2 PROTEIN IN

n g A	Tissues Lacking c-orbB-2 Protein
	a stage of the stage
	Politicatal oral muoosa ^{ce} Politicatal esophagus ^{ce}
	,
	Gk. rerulus ^{ez} Po. ratal Bowman's capsule ^{ez} Posmatal proximal tubule ^{ez}
	Pc tel colle cting ductes Pc triated renal palvis^{es} Pc triated fetal urater^{es} Uncorease
	Pu satic Işl eta^{sa}
٠.	Political traches ²² Political bronchioles ²²
	Pr - त्यांची alvooli ^{acaa} Pr - Julal brain ^{ac} Po - Julatal gangfion cells ^{ac} -
	Er Helium ^u

Lurtical cells⁶². ...! thymus⁶²

F: Lists⁴²
S th muscle cells⁴²
Centiac muscle cells⁴²

-hemical r. hose.

180 T.P. SINGLETON & DUC

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The data on case 3:0 should be interpreted. been studied, usually bodies. Studies using induce amplification:
been documented for

Activation of c-> -2

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carcinomas contained _ mif

epithelial malignanc: _ ;

carcinomas had c-erc

Activation of c->

TABLE & cerbB-2 AC.

gastric adenocarcino.

Turnor Ty

Overy—cardinoma, not : specified

Ovary—serous (pepillary)

Ovary—endometrioid ca

Ovary—mucinous cardino

Ovary—clear cell carcine
Ovary—mixed epithelial

Ovary—endometriold b. Ovary—mucinous bords

Overy—serous cystadens
Overy—mucinous cystad

Ovary-scierosing stror.

Ovary—fibrothecoma
Uterus—andometrial ads.

*Shown as number of case: given as supersoript. All pro and some or cases do not in the partished photover, are isted, with the immunity recipitation or ocen iden had in normal. Discrepancies regarding est in part, so differences in

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. perc.: i of **198) of** වර pe: 11 of **64) of**

TUMORE

c'erbE- mi N C ; pro	o-erbB-2 Protein Over- production
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subtypes (Table 281) of colorec study detected tion. One stud protein in colon

using Bouin's fi. sive carcinoma : erbB-2 protein

nomas (46 of 63 some of these "

TABLE 9. c-e/b8-

Esophagus-squa Stomach-carcine.

Stomach-carcino. Stomach-carcine:

Colorecturn-carci

Colon-villous ad Colon-tubulovillo. Colon-tubular ad. Colon-hyperplast Intestina-lelomyo. Hepatocellular card Hepatoblastoma Cholanglocarcinon Pancreas-adeno Pancreas—acinar Pancreas-dear c Pencreas-large c Pancreas-signet :

Shown as number of given as superscript. 2 mana

bTissues fixed in Bou

Conly cases with dis:

TABLE 10. o-erbB-

Tumor Type
Non-small cell carcin
Epidermold cardnors
Adenocarcinoma
Large cell carcinoma
Small cell cardinom.
Carcinoid tumor

Shown as aumber of c given as superscript. Al 2 mRNA.

does not indicate atic carcinomas a chemical reactive atic adenocarcine

c

Tables 10 th neoplasms. The Activation of e-e-non-small cell cafound e-erbB-2 p. noma had e-erbbtion of e-erbB-2 urinary bladder, cell carcinoma o

TABLE 11. C-OIDB

Tumor Ty:
Hematologic malign
Malignant lymphomi
Acute feukemia
Acute lymphoblastic
Acute myeloblastic i.
Chronic leukemia
Chronic lymphocytic
Chronic myelogenous
Myeloproliterative d.
Shown as number of a

given as superscript. Ali p

MON.

-	Protein Overproduction
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7 of 17).	rul cell carci-
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cerb8-2

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TABLE 12. c-eris . OFT : AND BONE -2 Dt licatio 1 0/89 227 0: 'nε 'n Shown as number of amber o studied; reference is given as superscript. I 2 proteiwhether the pr Enor kini.56 Thyroid carcinomas and rbB-2 mRNA. . ୯⊣• One abstract de lilq. in one of ten salivary gland pi Correlation of utc Very few studi rbBvation in nonmammary tumo d th B-2 amplification or overexpi s witi. cased survival, especially who ٠c T. 14. not report the stage, histologic ď , e. . asms. Another study of stages 11 ation between Ju on, ') decreased surviv t between survival and histolo rotein overpro-1-61 duction in 10 a vith decreased nrei : disease-free inte ·nd · immunohisto-TABLE 13. c-e TRACT o-erbB-2 Protein т. 31 Over-Tumor * " production Kidney-renal c Wilms' turnor Prostate-adeno 0/235 Urinary bladder-1/489 Shown as number / itudied; reference is given as superson, t...

184 T.P. SING STO	iD.		
TABLE 14. c-erbl A	<u>ন</u>	īUk.	
Tumor T 18		-2 mR. roduct	o-erbB-2 Protein Over- production
Sidn-malignant musine			0/10*
Skin, head and noc cell carcinoma)CT:	_	
Site not stated—sc carcinoma	ţi.	· -	-
Salivary gland—adenoc.	ጣሩ		_
Parolid gland—adenuid c carcinoma	:	-	0.141
Thyroid—anaplasti.		₹ <i>=</i>	=
Thyrold—papillary ir		'ev els)/ 5'	
Thyroid—aidenocar		_	
Thyroid—adenoma		vels)/2	_
Neuroblastoma			_
Meningioma			· _
Shown as number of comp	mp.		
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chemical reactiv	.î·t	Natabasa	
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SUMMARY			
Activation of the	2	⊳lificati	f c-erbB-2
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mately 20 perce.	şt	i a-eri	activation,
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